=> s 11

SAMPLE SEARCH INITIATED 14:06:52

SAMPLE SCREEN SEARCH COMPLETED -4 TO ITERATE

100.0% PROCESSED 4 ITERATIONS 1 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

> **COMPLETE** BATCH

PROJECTED ITERATIONS:

4 TO 200

PROJECTED ANSWERS:

1 TO 80

1 SEA SSS SAM L1

=> d scan

L2 1 ANSWERS REGISTRY COPYRIGHT 1997 ACS

Boronic acid, [3-methyl-1-[[1-oxo-3-phenyl-2-IN

[(pyrazinylcarbonyl)amino]propyl]amino]butyl]-, [S-(R*,S*)]- (9CI)

C19 H25 B N4 O4 MF

Absolute stereochemistry.

ALL ANSWERS HAVE BEEN SCANNED

=> s l1 full

FULL SEARCH INITIATED 14:08:09

FULL SCREEN SEARCH COMPLETED -64 TO ITERATE

100.0% PROCESSED 64 ITERATIONS

SEARCH TIME: 00.00.02

3 SEA SSS FUL L1 L3

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL **ENTRY** SESSION

3 ANSWERS

FULL ESTIMATED COST 112.88 113.03

FILE 'CAPLUS' ENTERED AT 14:09:49 ON 25 APR 1997 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 1997 AMERICAN CHEMICAL SOCIETY (ACS)

FILE COVERS 1967 - 25 Apr 1997 VOL 126 ISS 17 FILE LAST UPDATED: 25 Apr 1997 (970425/ED)

To help control your online searching costs, consider using the

HCAplus file when using the FSEARCH command or when conducting SmartSELECT searches with large numbers of terms.

Some chemical substances have deleted CAS Registry Numbers. To ensure that you are using the most current CAS Registry Number, and for a more complete search, start your CAS Registry Number search in the REGISTRY file. Then use the L-number answer set from REGISTRY as a search term in CAplus.

This file contains CAS Registry Numbers for easy and accurate substance identification.

```
=> s 13
L4
             1 L3
=> d bib abs hitstr
     ANSWER 1 OF 1 CAPLUS COPYRIGHT 1997 ACS
ΆN
     1996:466915 CAPLUS
DN
     125:143315
TI
     Boronic ester and acid compounds, synthesis and uses
     Adams, Julian; Ma, Yu-Ting; Stein, Ross; Baevsky, Matthew; Grenier,
IN
     Louis; Plamondon, Louis
     Proscript, Inc., USA
PA
     PCT Int. Appl., 144 pp.
SO
     CODEN: PIXXD2
                    960509
PI .
     WO 9613266 A1
DS
        AL, AM, AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES,
         FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG,
         SI, SĶ
     RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, DE, DK, ES, FR, GA, GB, GR,
         IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG
     WO 95-US14117 951027
PRAI US 94-330525 941028
     US 95-442581 950516
DT
     Patent
LΑ
     English
os
     MARPAT 125:143315
     Peptidyl boronic acids and esters PNR[B1R1X1]ACHR2X2CHR3BZ1Z2 [P =
AB
     aryl-, aralkyl-, heteroaryl-, or heteroarylalkylcarbonyl or
     -sulfonyl; B1 = N, CH; X1, X2 = CONH, CH(OH)CH2, COCH2; A = 0, 1, 2;
     R = H, alkyl; RR1 or RR2 (for A = 0) may form a ring; R1, R2, R3 =
     H, alkyl, cycloalkyl, aryl, etc.; Z1, Z2 = alkyl, hydroxy, alkoxy,
     aryloxy; Z1Z2 may form a moiety derived from a dihydroxy compd.] and
     their pharmaceutically acceptable salts were prepd. The rate of
     degrdn. of proteins of an animal can be reduced by contacting cells
     of the animal with these boronic compds. Thus, N-(4-
     morpholinecarbonyl)-.beta.-(1-naphthyl)-L-alanine-L-leucine boronic
     acid was prepd. by coupling (1S, 2S, 3R, 5S)-pinanediol leucine
     boronate trifluoroacetate salt with N-Boc-.beta.-(1-naphthyl)-L-
     alanine, followed by deprotection, acylation with
     4-morpholinecarbonyl chloride, and cleavage of the pinanediol
```

IT 179324-69-7 179324-85-7 179325-25-8
RL: BAC (Biological activity or effector, except adverse); BIOL

moiety.

Absolute stereochemistry.

RN 179324-85-7 CAPLUS

CN Boronic acid, [3-methyl-1-[[1-oxo-2-[(pyrazinylcarbonyl)amino]-3-(2-pyridinyl)propyl]amino]butyl]-, [S-(R*,S*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 179325-25-8 CAPLUS
CN Boronic acid, [1-[[3-(4-fluorophenyl)-1-oxo-2[(pyrazinylcarbonyl)amino]propyl]amino]-3-methylbutyl]-,
[S-(R*,S*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

=>

=> file beil

COST IN U.S. DOLLARS SINCE FILE TOTAL

FULL ESTIMATED COST ENTRY SESSION 7.81 120.84

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL

CA SUBSCRIBER PRICE ENTRY SESSION -0.48 -0.48

FILE 'BEILSTEIN' ENTERED AT 14:17:56 ON 25 APR 1997 COPYRIGHT (c) 1997 Beilstein Chemiedaten und Software GmbH, Beilstein Institut fuer Literatur der organischen Chemie

FILE LAST UPDATED: 07 APR 1997

FILE COVERS 1779 TO 1996.

*** CAS REGISTRY NUMBERS FOR 4,355,851 SUBSTANCES AVAILABLE ***

*** FILE CONTAINS 7,000,722 SUBSTANCES ***

- * PLEASE NOTE THAT THERE ARE NO FORMATS FREE OF COST.
- * SET NOTICE FEATURE: THE COST ESTIMATES CALCULATED FOR SET NOTICE
- * ARE BASED ON THE HIGHEST PRICE CATEGORY. THEREFORE; THESE
- * ESTIMATES MAY NOT REFLECT THE ACTUAL COSTS.
- * FOR PRICE INFORMATION SEE HELP COST *

=> d his

(FILE 'HOME' ENTERED AT 14:04:05 ON 25 APR 1997)

FILE 'REGISTRY' ENTERED AT 14:04:24 ON 25 APR 1997

L1 STRUCTURE UPLOADED

L2 1 S L1

L3 3 S L1 FULL

FILE 'CAPLUS' ENTERED AT 14:09:49 ON 25 APR 1997

L4 1 S L3

FILE 'BEILSTEIN' ENTERED AT 14:17:56 ON 25 APR 1997

=> s 11

SAMPLE SEARCH INITIATED 14:18:37

SAMPLE SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED 0 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.02

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 0 TO 0 PROJECTED ANSWERS: 0 TO 0

L5 0 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 14:19:50

FULL SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED 0 ITERATIONS . 0 ANSWERS

SEARCH TIME: 00.00.05

FULL ESTIMATED COST

L6 0 SEA SSS FUL L1

=> file marpat

COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION 0.00 120.84

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL

ENTRY SESSION

CA SUBSCRIBER PRICE 0.00 -0.48

FILE 'MARPAT' ENTERED AT 14:23:55 ON 25 APR 1997
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FILE CONTENT: 1988-PRESENT (VOL 104 ISS 14-VOL 126 ISS 16). (970418/ED)

MOST RECENT CITATIONS FOR PATENTS FROM FIVE MAJOR ISSUING AGENCIES (COVERAGE TO THESE DATES IS NOT COMPLETE):

US 5610305 11 MAR 1997

DE 19535340 27 MAR 1997

EP 764630 26 MAR 1997

JP 09077771 25 MAR 1997

WO 9709453 13 FEB 1997

Notice The first 1997 patent record appeared in MARPAT, with complete CA indexing and searchable Markush structure record, on 10 February 1997 -- US5591708 (970107), MARPAT 126:76542 -- just 5 weeks from issuance.

=> d his

(FILE 'HOME' ENTERED AT 14:04:05 ON 25 APR 1997)

FILE 'REGISTRY' ENTERED AT 14:04:24 ON 25 APR 1997 L1STRUCTURE UPLOADED L2 1 S L1 L3 3 S L1 FULL FILE 'CAPLUS' ENTERED AT 14:09:49 ON 25 APR 1997 L41 S L3 FILE 'BEILSTEIN' ENTERED AT 14:17:56 ON 25 APR 1997 L5 0 S L1 0 S L1 FULL L6 FILE 'MARPAT' ENTERED AT 14:23:55 ON 25 APR 1997 SAMPLE SEARCH INITIATED 14:25:06 SAMPLE SCREEN SEARCH COMPLETED -3 TO ITERATE 100.0% PROCESSED 3 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.11 FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE** PROJECTED ITERATIONS: 3 TO 164 PROJECTED ANSWERS: 0 TO 0 L7 0 SEA SSS SAM L1 => s 13 full FULL SEARCH INITIATED 14:25:36 FULL SCREEN SEARCH COMPLETED - 77 TO ITERATE 100.0% PROCESSED 77 ITERATIONS (2 INCOMPLETE) 3 ANSWERS SEARCH TIME: 00.00.18 Г8 3 SEA SSS FUL L1 => s 18 not 14 1 L4 L9 2 L8 NOT L4 => d bib abs qhit 1-2 ANSWER 1 OF 2 MARPAT COPYRIGHT 1997 ACS (ALL HITS ARE ITERATION INCOMPLETES) 125:143313 MARPAT ΑN ΤI Preparation of amidino and quanidino substituted peptide analogs as inhibitors of trypsin-like enzymes Lee, Sheng-lian O.; Carini, David John; Fevig, John Matthew; IN Kettner, Charles Adrian; Mantri, Padmaja; Feng, Zixia PΑ Du Pont Merck Pharmaceutical Company, USA PCT Int. Appl., 139 pp. SO CODEN: PIXXD2 PΙ WO 9612499 Al 960502 DS W: AU, CA, JP, MX, NZ. RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE WO 95-US13702 951024 ΑI

PRAI US 94-329039 941025 DT Patent LA English GI

$$Q^{1} = -(CH_{2})_{q}$$

$$Q^{1} = -(CH_{2})_{q}$$

$$(CH_{2})_{p}X$$

$$Q^{2} = -(CH_{2})_{q}$$

$$Q^{3} = -N$$

$$RNH - CH - B$$

$$(CH_{2})_{q}X$$

$$Me$$

$$Me$$

$$(CH_{2})_{q}X$$

$$Me$$

$$Me$$

$$Me$$

$$(CH_{2})_{q}X$$

AΒ Novel .alpha.-aminoacid and .alpha.-aminoboronic acid and corresponding peptide analogs of formula R3[A]nNR2CHR1E [E = BY1Y2, COR14, CO2R4, CONR15R16, COR4, COCO2R4; wherein Y1, Y2 = OH, F, (un) substituted NH2; or Y1Y2 = cyclic boron ester, cyclic boron amide, or cyclic boron amide-ester contg. 2-20 carbon atoms and optionally 1-3 heteroatoms selected from N, S, and O; R4 = H, C1-4 alkyl, aryl-C1-4 alkyl, C5-7 cycloalkyl; R14 = CF3, CHF2, CH2F, CH2Cl, CO2R4, CONR15R16, COR4, etc.; R15, R16 = H, C1-4 alkyl, aryl-C1-4 alkyl, C5-7 cycloalkyl, (un)substituted Ph; or NR15R16 = Q3; wherein W = single bond, O, S, SO, SO2, CH2, NR4, NCOR4; R1 = (un) substituted C1-12 alkyl, Q, Q1; wherein X = halo, cyano, NO2, CF3, NH2, NHC(:NH)H, NHC(:NH)NHOH, NHC(:NH)NHCN, etc.; Y = O, :NOH, :NNHCHO; p = 0-3; q = 0-4; R2 = H, (un)substituted C1-12 alkyl, cycloalkyl, Ph, naphthyl, or aryl-C1-4 alkyl; R3 = H, alkyl, aryl, alkylaryl, S(O)rR7, COR7, CO2R7, P(O)2OR7, or any other C1-20 NH2-blocking group; wherein R7 = H, C1-4 alkyl, (un)substituted Ph, naphthyl, or aryl-C1-4 alkyl; r = 0-2; A = amino acid residue orpeptide comprised of 2-20 amino acids residue; n = 0,1] and pharmaceutically acceptable salts thereof are prepd. These peptide analogs are useful for treating a physiol. disorder in a warm blooded animal catalyzed by trypsin-like enzymes, e.g. blood clotting, arterial thrombosis, myocardial infarction, inflammation, pancreatitis, and hereditary angioedema. Trypsin-like enzymes are a group of proteases which hydrolyze peptide bonds at basic residues liberating either a C-terminal arginyl or lysyl residue, among which are enzymes of the blood coagulation and fibrinolytic system required for hemostasis (e.g. factors II, X, VII, IX, kallikrein, tissue plasminogen activators, urokinase-like plasminogen activator, and plasmin), enzymes of the complement system, acrosin, and

pancreatic trypsin. Thus, Ac-D-Phe-Pro-OH was condensed with a boronic acid deriv. (I; R = H, X = Br) by a mixed anhydride procedure using iso-Bu chloroformate and N-methylmorpholine in CCl4 to give an intermediate I (R = Ac-D-Phe-Pro, X = Br), which was heated with Bu4NCN in MeCN at 90.degree. for 3 h to give the nitrile I (R = Ac-D-Phe-Pro, X = cyano). The latter nitrile was stirred with satd. methanolic HCl at 4.degree. overnight, concd., and redissolved in MeOH. NH3(g) was bubbled through the soln. for 1 h and the soln. was heated at 50.degree. for 3 h to give I [R = Ac-D-Phe-Pro, X = C(:NH)NH2]. This compd. in vitro inhibited thrombin with Ki of <500 nM.

MSTR 1A ITERATION INCOMPLETE

G1 = 7 / Hy<EC (3-6) Q (1) B (-5) N (-5) O (-3) S (0) OTHERQ (2-20) C, AN (1) B> / 12 / 51 / (SC 266)

G2 = OH / F / 10 / alkoxy<(1-8)>

G3 = H / alkyl<(1-4)> (SO G16) / cycloalkyl<(5-7)>
G4 = CF3 / 14 / 16 / 18 / 32 / 38 / 47 /
Hy<EC (1-4) Q (0-) N (0-) O (0-) S (0) OTHERQ, RC (1-3)>
(SO) / 207

G44=0 207

−NH---G20

-NH─G23

$$G15 = CN / 77$$

81 (O)-G21

G21 = H / alkyl<(1-4)> (SO G16) / Cb<EC (6-10) C, AR (1-), BD (ALL) N, RC (1-2), RS (1-2) E6 (0) OTHER> (SO) / OH / 83

RS (1-2) E6 (0) OTHER> (SO) / 81

0-----G22

G22 = alkyl<(1-4)> (SO G16) / Cb<EC (6-10) C, AR (1-), BD (ALL) N, RC (1-2),

```
RS (1-2) E6 (0) OTHER> (SO)
G23
        = H / alkyl < (1-4) > (SO G16) /
          Cb<EC (6-10) C, AR (1-), BD (ALL) N, RC (1-2),
          RS (1-2) E6 (0) OTHER> (SO) / OH / 95
95 (O)-G21
        = SH / 99 / 101 / 104 / CONH2 / 106 / 109 / 111 /
          CO2H / 172
                                                                     QН
. 99
                       G25-G26 C(O)·NH---G26 C(O)·G27
C(O)·O——G26
       = S / S(0) / SO2
G25
       = alkyl<(1-4)> (SO G16) / cycloalkyl<(5-7)>
       = H / alkyl < (1-4) > (SO G16) / cycloalkyl < (5-7) >
G27
       = SH / 120 / NH2
G29-G22
       = s / NH
G29
       = NH / 131 / 169
G30
      -G22 N—C(0)-G21
       = alkylene<EC (1-4) C, DC (0) M3>
= phenylene / cycloalkylene<(4-7)>
G31
G32
       = NULL / alkylene<EC (1-3) C, DC (0) M3> 
= R / (SC C(NH)NH2 / CH2NH2 / Br / CN / NH2 / OH /
G33
G34
          NHC(NH)NH2 / CO2Me)
G35
       = 0 / 187
N—
187
G36
       = OH / NHCHO
       = NH / 189
G37
```

-G18 N---= H / alkyl (SO G16) / Ph (SO) / naphthyl (SO) / SH / G38 191 / 193 / CHO / CO2H / 196 / 198 / 201 / (SC 346 / 350 / 359 / 361 / 373 / 382 / 391 / CO2CH2Ph / SO2Ph / CH2Ph / CH2CO2H) 191 G39-G40 196 C(O)·CH2—CH2—Ph 02S-350 С(O)-CH2—CH2—CO2H С(0)-CH2-CH2 с(о)-сн—сн́ G39 = S / S(O) / SO2 / C(O)= H / alkyl (SO G16) / Cb<EC (6-10) C, AR (1-), G40 BD (ALL) N, RC (1-2), RS (1-2) E6 (0) OTHER> (SO) G41 = OH / 205205 -G42 G42 = alkyl (SO G16) / Ph (SO) / naphthyl (SO)

= R<TX "peptide residue of 1-20 amino acids"> /

(SC 209-1 211-3 / 217-1 218-3 / 221-1 223-3 / 261-1 258-3 / 298-1 295-3 / 309-1 314-3 / 332-1 323-3 / 341-1 338-3)

G43

G44 = Hy < EC (1-4) Q (0-) N (0-) O (0-) S (0) OTHERQ,RC (1-3) > (SO)

G45 = Me / CH2CH2CH2NHC(NH)NH2 / CH2CONH2 / CH2CO2H /
CH2SH / CH2CH2CONH2 / CH2CH2CO2H / H / 224 / 231 / Bu-s /
Bu-i / CH2CH2CH2NH2 / CH2CH2SMe / CH2CH2CH2NH2 / CH2Ph /
240 / CH2OH / CH(OH)Me / 242 / CH2C6H4OH-p / Pr-i

= phenylene / cycloalkylene<(4-7)>

```
G47
      = Ph / thiazolyl / 2-pyridyl / 3-pyridyl / 2-thienyl
      = CH2CH=CH2 / Me / Et
         and pharmaceutically acceptable salts
DER:
MPL:
        claim 1
         substitution is restricted
NTE:
NTE:
         alkyl groups in G18 may contain heteroatom interruptions
         210,216,222 - D,L; 260,297 - D
STE:
    ANSWER 2 OF 2 MARPAT COPYRIGHT 1997 ACS
(ALL HITS ARE ITERATION INCOMPLETES)
AN
     123:957 MARPAT
ΤI
     Electrophilic peptide analogs as inhibitors of trypsin-like enzymes
TN
     Galemmo, Robert Anthony, Jr.; Abelman, Matthew Mark; Amparo, Eugene
     Cruz; Fevig, John Matthew; Knabb, Robert Madara; Miller, William
     Henry; Pacofsky, Gregory James; Weber, Patricia Carol
PA
     Du Pont Merck Pharmaceutical Co., USA
so
     PCT Int. Appl., 307 pp.
     CODEN: PIXXD2
PΙ
    WO 9509634 A1
                    950413
    W: AU, CA, CZ, FI, HU, JP, KR, NO, NZ, PL, SK
DS
     RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
AΤ
     WO 94-US11280 941006
PRAI US 93-133251 931007
     US 93-139445 931020
DT
     Patent
LΑ
     English
AB
     Electrophilic dipeptide analogs R3R11NCR4R5C(O)NHCHR1A [R1 = ZX; Z =
     C1-12 alkyl or alkenyl, (CH2) qC6H4(CH2)p; p = 0-3; q = 0-4; X = 0-4
     halo, CN, NO2, CF3, NH2, etc.; R3 = C(0)Y; Y = aryl, aralkyl,
     heterocyclyl, heterocyclylalkyl, cycloalkylalkyl, adamantylalkyl,
     etc.; R4, R5 = H, C1-4 alkyl, (C1-4 alkyl)aryl, C5-7 cycloalkyl; R11
     = C1-4 alkyl, C3-6 cycloalkyl, alkoxy, NH2, (di)alkylamino, aryl,
    heterocyclyl, etc.; A = B(OH)2, BF2, cyclic B ester or amide,
     C(O)CF3, C(O)C(O)NH2, CH(OH)(CH2F), etc.] in which an electrophilic
     deriv. of an .alpha.-amino acid is conjugated to an
     N, N-disubstituted .alpha.-amino acid are prepd. as inhibitors of
     trypsinlike serine proteases for use as antithrombotics. Thus,
     N-hydrocinnamoy1-N-(2,2-dimethy1-2-phenylethy1)glycylborolysine-HCl
     (I) was prepd. by condensation of H2NCH2C(O)OEt.HCl with PhCMe2CHO
     (prepn. given) followed by PhCH2CH2C(0)Cl and Br(CH2)4CH(NH2)B(OH)2
     pinanediol ester and transesterification with PhB(OH)2. I showed Ki
     <500 nM for thrombin, Factor Xa, and Factor VIIa using synthetic
     chromogenic substrates and IC50 <500 nM for thrombin time.
```

MSTR 1C ITERATION INCOMPLETE

= Ak<EC (1-12) C, BD (0-) D (0) T> / phenylene / $99-6\ 100-9$ / $101-6\ 102-9$ / $10-6\ 12-9$ G1 G2—G3—G4 10 12 99 100 G2-G34 101 102 G2 = alkylene<(1-4)>G3 = phenylene = alkylene<(1-3)>G4 = CN / CF3 / 20 / 23 23 (O)-G9 = NH2 / alkylamino<(1-4)> / NHCHO / G8 alkylcarbonylamino<(1-4)> = NH2 / alkylamino<(1-4)> / OH / alkoxy<(1-4)> = 34 / 48 / 64 / 103 / 145 / 110 / 257 / 262 / 267 G9 G10 C(0)-G11—G14 C(0)-G21-G22 C(0)-G24 C(0)G45 C(0)G44 G31 G31 G38 G39 N2

= NULL / alkylene (SO (-2) G12) / alkenylene<(2-5) > /

39-34 40-36 / 45-34 47-36

C(0) G45 C(0) G44 G31

G11

```
G15-G18 G18-G19-G20
G12
       = alkoxy<(1-4)>/ aryl<RC (1-3),
         RS (0-1) E5 (1-2) E6 (0) OTHER> (SO) /
         Hy<EC (1-2) Q (0-) N (0-) O (0-) S (0) OTHERQ, RC (1-2),
         RS (-1) E5 (-2) E6 (0) OTHER> (SO) /
         aryloxy<RC (1-3), RS (0-1) E5 (1-2) E6 (0) OTHER> (SO) / 37
C (0)-G13
G13
       = OH / alkoxy<(1-4)> (SO aryl<RC (1-3),
         RS (0-1) E5 (1-2) E6 (0) OTHER> (SO)) / cycloalkyloxy<(5-7)>
       = aryl < RC (1-3), RS (0-1) E5 (1-2) E6 (0) OTHER>
G14
         (SO) / Hy < EC (1-3) Q (0-) N (0-) O (0-) S (0) OTHERQ,
         RC (1-2), RS (-1) E5 (-2) E6 (0) OTHER> (SO)
       = 0 / S / S(0) / SO2 / 41
G15
N---G16
G16
       = H / alkyl < (1-4) > (SO aryl < RC (1-3),
         RS (0-1) E5 (1-2) E6 (0) OTHER> (SO)) / cycloalkyl<(5-7)> /
C(0)-G17
G17
       = H / alkyl < (1-4) > (SO aryl < RC (1-3),
         RS (0-1) E5 (1-2) E6 (0) OTHER> (SO)) / cycloalkyl<(5-7)>
       = CH2 (SO)
G18
       = 0 / S / S(0) / S02
G19
G20
       = (0-2) CH2
       = (1-3) CH2
G21
G22
       = adamantyl / cycloalkyl<(5-7)> / 51
G42-G23
G23
       = cycloalkyl<(5-7)>
       = Ph (SR) / 58 / 183 / 192 / 204 / 234 / 248
G24
```

$$G^{25}$$
 G^{25}
 G^{25}

G25 = H / R G26 = S / S(O) / SO2 G27 = 73 / Hy<EC (3-6) Q (1) B (0-) N (0-) O (0-3) S (0) OTHERQ (2-20) C> / 76 / 78

```
= OH / F / NH2 (SO) / alkoxy<(1-8)>
G28
      = R / (EX H)
G29
      = R / (EX H)
G30
      = H / R
G31
G32
       = alkyl<(1-4)> (SO aryl<RC (1-3),
         RS (0-1) E5 (1-2) E6 (0) OTHER> (SO)) / cycloalkyl<(3-6)> /
         OH (SO) / NH2 (SO) / CONH2 (SO) /
         aryl<RC (1-3), RS (0-1) E5 (1-2) E6 (-1) E7 (0) OTHER> (SO) /
         Hy<EC (1-3) Q (0-) N (0-) O (0-) S (0) OTHERQ, RC (1-2),
         RS (-1) E5 (-2) E6 (0) OTHER> (SO) /
         alkyl<(1-4)> (SR Hy<EC (1-3) Q (0-) N (0-) O (0-) S (0)
         OTHERQ, RC (1-2), RS (-1) E5 (-2) E6 (0) OTHER> (SO)) /
         alkyl<(1-4)> (SR CO2H (SO)) / 273 / 276 / 283 / 288
```

G33 = phenylene G34 = phenylene G35 = naphthyl (SO)

= 115 / 149 / 163 / 210 / 218 G36

G37 = (0-3) CH2 G38 = Ph (SO) G39 = (0-4) CH2 G40 = (0-1) CH2 G41 = (0-2) CH2 = 0 / S / S(0) / S02 / NH (S0)G42 = NH2 (SO) G43 G44 = 0 / NH= Ak < EC (3-) C, BD (ALL) SE > (SO) G45 = H / alkyl < (1-4) > (SO aryl < RC (1-3),G46 RS (0-1) E5 (1-2) E6 (0) OTHER> (SO)) / cycloalkyl<(5-7)> G50 = alkylene G32+G46= CH2CH2 or pharmaceutically acceptable salts, hydrates or prodrugs DER:

MPL: claim 1

NTE: additional ring formation allowed

substitution is restricted => => => s 19/com LIMIT NOT VALID FOR L9 This qualification can be applied only to a structure answer set L-number. => => d his (FILE 'HOME' ENTERED AT 14:04:05 ON 25 APR 1997) FILE 'REGISTRY' ENTERED AT 14:04:24 ON 25 APR 1997 STRUCTURE UPLOADED L1L2 1 S L1 L3 3 S L1 FULL FILE 'CAPLUS' ENTERED AT 14:09:49 ON 25 APR 1997 L41 S L3 FILE 'BEILSTEIN' ENTERED AT 14:17:56 ON 25 APR 1997 L50 S L1 0 S L1 FULL L6 FILE 'MARPAT' ENTERED AT 14:23:55 ON 25 APR 1997 0 S L3 L7 L8 3 S L3 FULL 2 S L8 NOT L4 L9 => s 18/com1 L8/COM L10 => s 110 not 14 1 L4 L11 0 L10 NOT L4 => => logoff ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF LOGOFF? (Y)/N/HOLD:y COST IN U.S. DOLLARS SINCE FILE TOTAL SESSION ENTRY FULL ESTIMATED COST 54.09 174.93 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -0.92 -1.40

STN INTERNATIONAL LOGOFF AT 14:35:48 ON 25 APR 1997